Nipah Viral Virulence may be Due to Resistant Strains Amplified by Cross-Species Transmission between Animals and Humans

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Abstract

Nipah virus (NiV) is a newly emerging devastating viral infection that causes severe illness and loss of both animals and humans lives. Genetic resistant strains of NiV amplify the virulence through cross transmission between animals and humans. It is very dangerous and engulfs lives of people within short notice. The dearth of specific chemotherapeutics and vaccination lead to the study of dynamic viral strains. The present study represents an exploration of different NiV targets for the development of potential antiviral compounds. The manuscript describes more about the current scenario of death statistics, various NiV targets, available treatment options and research endeavors to discover a treatment strategy.

Keywords: Nipah virus (NiV); Virulence; Cross-species transmission between animals and humans; Drug targets; Genetically resistant strains.

Introduction

Past pandemic and epidemic of NiV

Nipah virus is one of the ribonucleic acid viruses belong to the family members of Paramyxoviridae, genus Henipavirus. NiV was first originated from Sungai Nipah, a village in the Malaysian Peninsula where pig farmers became ill due to respiratory tract infection followed by encephalitis. NiV attacked pig farmers or abattoir workers which were first reported in September 1998 to April 1999 in Malaysia and then spread to neighbor Singapore by March 1999 [1,2] and in India and Bangladesh in 2001 [3]. Many 105 people died in Malaysia in April 1999, and one fellow was died in Singapore in the same year [1]. In 2001, a number of 45 persons were died followed by the death of 5 persons in 2007 in India whereas from 2001-2013 a number of 221 infected out of which 171 were being died in Bangladesh [4]. The morbidity and mortality statistics are given in Table 1.

The people in contact with the infected pigs get an infection. Infected people migrated in various countries and further infection speeded to Cambodia [5], Thailand [6], Indonesia [7], Madagascar in Southern Africa [8] and Ghana in West Africa [9,10].

Infected people initially develop symptoms including fever, headaches, myalgia (muscle pain), vomiting and sore throat. This can be followed by dizziness, drowsiness, altered consciousness, and neurological signs that indicate acute encephalitis.

Recent outbreaks in India in May 2018

A recent outbreak at Kerala, India on May 31, 2018, recorded 18 deaths due to suspected Nipah virus infection. The death of three members of a family in Changanthothu panchayath was confirmed as due to Nipah virus in the tests conducted at National Virology Institute, Pune. The latest victim suspected to have died due to Nipah infection is a lady of 31 years old, a nurse of the Perambra Taluk Hospital. She was taking care of the patients and was later confirmed to have Nipah infection at the Perambra hospital [11]. Scientists are worried due to lack of specific treatment to eradicate NiV.

Mode of treatments

In vitro study confirmed anti-Nipah viral activity of ribavirin has been shown effective in vitro tests, but has not yet been proven effective in humans and has to be passed in different clinical trials [12]. Passive immunization using a human monoclonal antibody that targets the Nipah G glycoprotein has been evaluated in the ferret model as post-exposure prophylaxis [13,14]. The anti-malarial drug chloroquine has been used in people on a compassionate use basis in Australia and was in pre-clinical development in 2013 [13]. Favipiravir (T-705) is a purine analog antiviral approved for use in Japan against emerging influenza strains, and several phases 2 and 3 clinical trials are ongoing in the United States and Europe. Favipiravir has demonstrated efficacy against a broad spectrum of RNA viruses, including members of the Paramyxoviridae, Filoviridae, Arenaviridae families, and the Bunyaviridae order. The scarcity of specific chemotherapeutics emphasizes the priority of exploring different targets of NiV. Dawes et al. reported that favipiravir has potent in vitro antiviral activity against Nipah and Hendra virus replication and transcription at micromolar concentrations [15]. Recently, the adenosine nucleoside analog GS-441524, and its monophosphate prodrug GS-5734 were demonstrated to have in vitro antiviral activity against NiV and HeV with EC50 values between 0.49 to 1 μM and 0.032 to 0.055 μM, respectively [16].

Quest of different drug targets

NiV has the Nipah virus is a type of RNA virus in the genus Henipavirus.
having a morphology similar to other paramyxoviruses with pleomorphic structure, herringbone nucleocapsids, and 10 nm long surface protein spikes on the envelope (Figure 1). Ribonucleic acid of NiV is well settled within the lipid bilayer coat containing six genes such as N, P, M, F, G and L responsible for producing nucleoprotein, phosphoprotein, matrix, fusion, glycoprotein and large RNA polymerase [17]. Complex formation between nucleoprotein and phosphoprotein component viral chaperone is responsible for the viral replication. Such complex is called as N0–P core complex, the crystal structure of which was solved by Yabukarksi and colleagues [18].

The NiV G and F proteins mediate the entry of genome, attachment, and fusion with the host receptor cells [19]. Ephrin-B2 (EFNB2) was recognized as a human host cellular receptor for the attachment of NiV glycoproteins. Such attachment is mediated by the sialic acid interactions having phenylalanine side chain on EFNB2 that fits snugly into a hydrophobic pocket on the viral protein [20]. The fusion protein, an enveloped glycoprotein essential for viral entry, belongs to the family of class I fusion proteins and is characterized by the presence of two heptad repeat (HR) regions, HR1 and HR2. These two regions associate to form a fusion-active hairpin conformation that juxtaposes the viral and cellular membranes to facilitate membrane fusion and enable subsequent viral entry. The Hendra and Nipah virus fusion core proteins were crystallized by Lou and co-workers [21]. Matrix protein is situated beneath the lipid bilayer and helps to connect with the viral RNA that is accountable for the formation of its virions particles responsible for the virulence effect causing gene expression and immune evasion.

Mode of spreading of NiV infection

Fruit bats have an affinity to taste the pulpy fruits such as bananas, mangos, dates, avocados, guava, papaya, figs and wild dates bearing thin peel as well as some vegetables like a banana flower and bean. NiV is being transmitted by the bats that feed on or touches. Figure 2 denotes fruits bat eating guava (A), banana (B), date palm (C), papaya (D), mango (E), figs (F), the flower of banana (G) and beans (H). Humans and animals eat the infected fruits and transmission of the disease occurs.

Fruit bats live at the date and palm trees. Date palm sap is so delicious and collected by the villagers who climb the tree and come in touch with the virus. Date palm sap can be infected by the saliva and excretory materials secreted by the fruit bats containing NiV. Figure 3 indicates the villager collects date palm sap (A), palm sap (B) in a clay pot (C), and fruit bats attacking stem of the date tree (D) and sap (E).

Luby et al. [22] suggested that NiV was transmitted from fruit bats to persons through drinking fresh date palm sap. Date palm sap occasionally contaminated with NiV-infected bat urine or saliva that contains a sufficient dose of NiV to be fatal to humans. In India, in a bat sample survey, NiV RNA was detected in a liver homogenate of fruit bat.
bats captured in Myanaguri, West Bengal [23].

Expert's Opinion

Deforestation leads to the destruction of natural habitat followed by global warming and ecological imbalance which may cause homeless to the fruits bats, also called as flying foxes, who moved to the neighbor states and countries. Fruits bats of the genus Pteropus close in contact with the pigs came into the picture as a carrier of NiV. The natural reservoir of fruit bats is NiV and the carrier is animals such as pigs and horses. NiV is transmitted into the healthy humans in close contact with the infected pigs and horses or upon the consumption of contaminated fruits and date palm sat. Then viral strains cross-transmission between humans and these animals cause genetically resistant strains which are so dreadful (Figure 4). When these animals sneeze, cough, secrete tears and passing stools, the resistant viral strains may come out through their nasopharyngeal secretions, tears, and intestinal diarrheal materials and can infect the healthy humans in close contact with this contaminated environment having viral genetic mutant drift particles. The virus is transmitted from the carrier (horses and pigs) to humans through consumption of contaminated foods or upon contact with the respiratory droplets or body fluids of infected animals. This appears to be a case of cross-species transmission (CST) [24] of a same strain of virus between different organisms which produces resistant strains rather than genetic re-assortment. Cross-species transmission causes amplification to the gene mutations leading to generate highly dangerous resistant NiV strain which is responsible for the NiV virulence that may overcame the host immune defense mechanism. It has been assumed that a Keralian nurse died due to infection with the dangerous resistant NiV strain. Further genomic scientists can analyze the genome of dangerous viral strains which would be helpful for the development of anti-Nipah viral drug and vaccination. More attention has to be paid to the discovery of anti-Nipah viral drug and vaccine. Existing antiviral strain is a major target to explore its genome sequences for the design and discovery of anti-Nipah viral drug and vaccine. Existing antiviral drugs could be repurposed to test against NiV.

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References